House	Amendment NO
Offered By	
AMEND Senate Bill No. 717, Pag following:	ge 1, Section A, Line 2, by inserting immediately after said line the
"191.480. 1. For purposes	of this section, the following terms shall mean:
(1) "Eligible patient", a pe	erson who meets all of the following:
(a) Has a terminal illness;	
(b) Has considered all other	er treatment options currently approved by the United States Food
and Drug Administration and all re	elevant clinical trials conducted in this state;
(c) Has received a prescrip	otion or recommendation from the person's physician for an
investigational drug, biological pro	oduct, or device;
(d) Has given written infor	rmed consent for the use of the investigational drug, biological
product, or device or, if the patient	is a minor or lacks the mental capacity to provide informed
consent, a parent or legal guardian	has given written informed consent on the patient's behalf; and
(e) Has documentation from	m the person's physician that the person has met the requirements
of this subdivision;	
(2) "Investigational drug, b	biological product, or device", a drug, biological product, or
device that has successfully comple	eted phase one of a clinical trial but has not been approved for
general use by the United States Fo	ood and Drug Administration and remains under investigation in a
clinical trial. The term does not in	clude Schedule I controlled substances;
(3) "Terminal illness", a di	isease that without life-sustaining procedures will result in death in
the near future or a state of perman	nent unconsciousness from which recovery is unlikely.
2. A manufacturer of an in	vestigational drug, biological product, or device may make
available the manufacturer's invest	tigational drug, biological product, or device to eligible patients
	es not require that a manufacturer make available an
	oduct, or device to an eligible patient. A manufacturer may:
(1) Provide an investigation	onal drug, biological product, or device to an eligible patient
without receiving compensation; o	
	ient to pay the costs of or associated with the manufacture of the
investigational drug, biological pro	<del></del>
	quire a health care insurer to provide coverage for the cost of any
investigational drug, biological pro	oduct, or device. A health care insurer may provide coverage for
Action Taken	Date

an investigational drug, biological product, or device.

- 4. Notwithstanding any other provision of law to the contrary, no state agency or regulatory board shall revoke, fail to renew, or take any other action against a physician's license issued under chapter 334 based solely on the physician's recommendation to an eligible patient regarding prescription for or treatment with an investigational drug, biological product, or device.
- 5. Any official, employee, or agent of this state who blocks or attempts to block access of an eligible patient to an investigational drug, biological product, or device is guilty of a class A misdemeanor.
- 6. If a provision of this section or its application to any person or circumstance is held invalid, the invalidity does not affect other provisions or applications of this section that can be given effect without the invalid provision or application, and to this end the provisions of this section are severable."; and

- 14 Further amend said bill by amending the title, enacting clause, and intersectional references
- 15 accordingly.